STM-Structure Scarch

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ANSWER 1 OF 7 CAPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER:

2003:913040 CAPLUS

DOCUMENT NUMBER:

139:375018

TITLE:

Combinations containing proton pump inhibitors for the

treatment of airway disorders

INVENTOR(S):

Hanauer, Guido; Kromer, Wolfgang; Postius, Stefan;

Simon, Wolfgang-Alexander

PATENT ASSIGNEE(S):

Altana Pharma A.-G., Germany

SOURCE:

PCT Int. Appl., 21 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.					KIND DATE				APPLICATION NO.						DATE		
				A3	2 20031120 3 20040401										20030503		
	RW:	VN, AT, IT,	YU, BE, LU,	ZA, BG, MC,	ZW, CH,	LV, AM, CY,	CA, MA, AZ, CZ, RO,	MK, BY, DE,	MX, KG, DK,	NO, KZ, EE,	NZ, MD, ES,	PH, RU,	PL, TJ.	SG, TM	TN,	UA,	US,
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AΒ inhibitor is, e.g., Soraprazan or its salt, and the airway therapeutic is, e.g., Ciclesonide.

ΙT 362525-74-4

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (oral combination of reversible proton pump inhibitors and airway therapeutics for treatment of airway disorders)

362525-74-4 CAPLUS RN

7H-Imidazo[1,2-a]pyrano[2,3-c]pyridin-8-ol, 8,9-dihydro-7-(2-CN methoxyethoxy)-2,3-dimethyl-9-phenyl-, (7R,8R,9R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L4ANSWER 2 OF 7 CAPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER:

2003:875288 CAPLUS

DOCUMENT NUMBER:

139:364931

TITLE:

Preparation of nitrosated tricyclic imidazopyridine derivatives as gastric secretion-inhibitor and

anti-inflammatory and antibacterial agents INVENTOR(S):

Buhr, Wilm; Senn-Bilfinger, Joerg; Zimmermann, Peter

Jan

PATENT ASSIGNEE(S):

SOURCE:

Altana Pharma Ag, Germany PCT Int. Appl., 62 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

Ι

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT	PATENT NO.						KIND DATE		APPLICATION NO.					DATE		
WO 2003				WO 2003-EP4134						20030422						
W:	ΑE,	AL,	ΑU,	BA,	BR,	CA,	CN,	CO,	CU,	DZ,	EC,	GE,	HR,	ID.	IL.	IN.
	IS,	JP,	KR,	LT,	LV,	MA,	MK,	MX,	NO,	NZ,	PH,	PL.	SG.	TN.	UA.	US
	VN,	YU,	ZA,	ZW,	ΑM,	ΑZ,	BY,	KG,	KZ,	MD,	RU,	ТJ,	TM		•	-
RW:	ΑT,	ΒE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FΙ,	FR,	GB,	GR,	HU,	IE.
	IT,	LU,	MC,	NL,	PT,	RO,	SE,	SI,	SK,	TR			·	•	•	
PRIORITY APP	. :			·	•			002-	9104		7	A 20	00204	121		
OTHER SOURCE	MARPAT 139:364931										7020-	124				

OTHER

GI

The invention relates to nitrosated tricyclic imidazopyridines (e.g. AΒ 7,8,9,10-tetrahydroimidazo[1,2-h][1,7]naphthyridine) of formula (I) [R1 = H, C1-4 alkyl, C3-7 cycloalkyl, C3-7 cycloalkyl-C1-4 alkyl, C1-4 alkoxy, C1-4 alkoxy-C1-4 alkyl, C1-4 alkoxycarbonyl, C2-4 alkenyl, C2-4 alkynyl, fluoro-C1-4 alkyl, hydroxy-C1-4 alkyl; R2 = H, C1-4 alkyl, aryl, C3-7 cycloalkyl, C3-7 cycloalkyl-C1-4 alkyl, C1-4 alkoxycarbonyl, hydroxy-C1-4 alkyl, halogen, C2-4 alkenyl, C2-4 alkynyl, fluoro-C1-4 alkyl, cyanomethyl, etc.; R3a, R3b = H, halogen, fluoro-C1-4 alkyl, C1-4 alkyl, C2-4 alkenyl, C2-4 alkynyl, CO2H, -CO-C1-4 alkoxy, hydroxy-C1-4 alkyl, C1-4 alkoxy-C1-4 alkyl, C1-4 alkoxy-C1-4 alkoxy-C1-4 alkyl, fluoro-C1-4 alkoxy-C1-4 alkyl, (un)substituted CONH2; one of R4a and R4b or one of R5a and R5b = H, C1-7 alkyl, C2-7 alkenyl, Ph or phenyl-C1-4 alkyl and the other = HO, C1-4 alkoxy, oxo-substituted C1-4 alkoxy, C3-7 cycloalkoxy, C3-7 cycloalkyl-C1-4 alkoxy, hydroxy-C1-4 alkoxy, C1-4 alkoxy-C1-4 alkoxy, C1-4 alkoxy-C1-4 alkoxy-C1-4 alkoxy, C3-7 cycloalkoxy-C1-4 alkoxy, C3-7 cycloalkyl-C1-4 alkoxy-C1-4 alkoxy, C1-4 alkylcarbonyloxy, wholly or mainly halogen-substituted C1-4 alkoxy, etc. or in which R4a and R4b or R5a and R5b together are O (oxygen) or are C1-7 alkylidene; Arom = (un) substituted mono- or bicyclic aromatic radical; X = 0 or NH]. disclosed is the use of the compds. I for the prevention and treatment of gastrointestinal illnesses. These compds. are acid pump antagonists (APAs) with less side effects than known APAs and have an antibacterial activity against Helicobacter bacteria with less side effects than known compds. with such activity and NO (nitric oxide) releasing activity, in

which the effect against Helicobacter bacteria is synergistically enhanced on account of the gastric acid inhibiting activity of these compds. They exhibit a marked inhibition of gastric secretion and an excellent gastric and intestinal protective action in warm-blooded animals, in particular humans. Due to gastric and intestinal protection, they are useful for the prevention and treatment of gastrointestinal diseases, in particular of gastrointestinal inflammatory diseases and lesions (e.g. gastric ulcer, peptic ulcer, including peptic ulcer bleeding, duodenal ulcer, gastritis, hyperacidic or medicament-related functional dyspepsia), which can be caused, for example, by microorganisms (e.g. Helicobacter pylori), bacterial toxins, medicaments (e.g. certain antiinflammatories and antirheumatics, such as NSAIDs and COX-inhibitors), chems. (e.g. ethanol), gastric acid or stress situations.

IT 620631-28-9P

CN

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(intermediate; preparation of nitrosated tricyclic imidazopyridine derivs. as gastric secretion-inhibitor and anti-inflammatory and antibacterial agents for prevention and treatment of gastrointestinal diseases)

RN 620631-28-9 CAPLUS

Pentanoic acid, 5-bromo-, (7R,8S,9R)-8,9-dihydro-7-(2-methoxyethoxy)-2,3-dimethyl-9-phenyl-7H-imidazo[1,2-a]pyrano[2,3-c]pyridin-8-yl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

IT 620631-26-7P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of nitrosated tricyclic imidazopyridine derivs. as gastric secretion-inhibitor and anti-inflammatory and antibacterial agents for prevention and treatment of gastrointestinal diseases)

RN 620631-26-7 CAPLUS

CN Pentanoic acid, 5-(nitrooxy)-, (7R,8S,9R)-8,9-dihydro-7-(2-methoxyethoxy)-2,3-dimethyl-9-phenyl-7H-imidazo[1,2-a]pyrano[2,3-c]pyridin-8-yl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

IT 362605-90-1

RL: RCT (Reactant); RACT (Reactant or reagent) (reactant; preparation of nitrosated tricyclic imidazopyridine derivs. as gastric secretion-inhibitor and anti-inflammatory and antibacterial agents for prevention and treatment of gastrointestinal diseases)

362605-90-1 CAPLUS RN

7H-Imidazo[1,2-a]pyrano[2,3-c]pyridin-8-ol, 8,9-dihydro-7-(2-CN methoxyethoxy)-2,3-dimethyl-9-phenyl-, (7R,8S,9R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 3 OF 7 CAPLUS COPYRIGHT 2004 ACS on STN

3

ACCESSION NUMBER: 2003:417606 CAPLUS

DOCUMENT NUMBER: 139:946

TITLE: Reversible proton pump inhibitors for the treatment of

airway disorders

INVENTOR(S): Senn-Bilfinger, Joerg; Kassel, Gerd; Hanauer, Guido;

Buhr, Wilm; Simon, Wolfgang-Alexander

PATENT ASSIGNEE(S): Altana Pharma A.-G., Germany

SOURCE: PCT Int. Appl., 18 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

REFERENCE COUNT:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE				
WO 2003043614 WO 2003043614	A2 A3	20040311	20040311					
IN, IS, JP,	KR, LT	, LV, MA, MK,	CU, DZ, EC, GE, HR, MX, NO, NZ, PH, PL, AZ, BY, KG, KZ, MD,	RO. SG. ST.				

RN 533903-14-9 CAPLUS

CN 7H-Imidazo[1,2-a]pyrano[2,3-c]pyridin-8-ol, 7-butoxy-8,9-dihydro-2,3-dimethyl-9-phenyl-, (7R,8R,9R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 533903-15-0 CAPLUS

CN 7H-Imidazo[1,2-a]pyrano[2,3-c]pyridin-8-ol, 7-butoxy-8,9-dihydro-2,3-dimethyl-9-phenyl-, (7S,8R,9R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L4 ANSWER 4 OF 7 CAPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER:

2001:730749 CAPLUS

DOCUMENT NUMBER:

135:272986

TITLE:

Preparation of imidazopyridine prodrugs for prevention

and treatment of gastrointestinal diseases

INVENTOR(S):

Simon, Wolfgang-Alexander; Postius, Stefan; Huber, Reinhard; Kromer, Wolfgang; Senn-Bilfinger, Joerg;

Buhr, Wilm

PATENT ASSIGNEE(S):

Byk Gulden Lomberg Chemische Fabrik G.m.b.H., Germany

SOURCE:

PCT Int. Appl., 59 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

Engli

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE ______ ----------WO 2001072756 A120011004 WO 2001-EP3514 20010328 W: AE, AL, AU, BA, BG, BR, CA, CN, CO, CZ, EE, GE, HR, HU, ID, IL, IN, JP, KR, LT, LV, MK, MX, NO, NZ, PL, RO, SG, SI, SK, UA, US, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR AU 2001060166 A5 20011008 AU 2001-60166 20010328 EP 1313740 A1 20030528 EP 2001-933769 20010328 AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR BR 2001009483 Α 20030610 BR 2001-9483 20010328 JP 2003528878 T2 20030930 JP 2001-570665 20010328 NO 2002004662 Α 20020927 NO 2002-4662 20020927 US 2003125327 Α1 20030703 US 2002-182619 20021001 PRIORITY APPLN. INFO.: A 20000329 Sell EP 2000-106695

WO 2001-EP3514

W 20010328

OTHER SOURCE(S):

MARPAT 135:272986

GΙ

AB Imidazopyridines, such as I [R4, R5 = OH, alkoxy, alkylcarbonyloxy, carbamoyloxy, alkyloxycarbonyloxy, etc.], were prepared for pharmaceutical use as prodrugs for the treatment of gastrointestinal disorders, such as gastrointestinal inflammatory diseases and lesions and gastric acid related diseases. Thus, imidazopyridine II [R4 = O(CH2)2OMe, R5 = COMe] was prepared via O-alkylation of the corresponding diol II (R4 = R5 = OH) with MeO(CH2)2OH followed by acetylation with acetic anhydride. The prepared imidazopyridines were tested for their inhibition of stomach acid secretion of perfused rat stomach stimulated by pentagastrin.

IT 362525-48-2P 362525-50-6P 362525-52-8P 362525-54-0P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of imidazopyridine prodrugs for prevention and treatment of gastrointestinal diseases)

RN 362525-48-2 CAPLUS
CN 7H-Imidazo[1,2-alpy

7H-Imidazo[1,2-a]pyrano[2,3-c]pyridin-8-ol, 8,9-dihydro-7-(2-

REFERENCE COUNT:

THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS 4 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 5 OF 7 CAPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER:

2001:730748 CAPLUS

DOCUMENT NUMBER:

135:272963

TITLE:

Preparation of pyrano[2,3-c]imidazo[-1,2-a]pyridine derivatives for the treatment of gastrointestinal

disorders

INVENTOR(S):

Simon, Wolfgang-alexander; Postius, Stefan; Kromer,

Wolfgang; Senn-bilfinger, Joerg; Buhr, Wilm

PATENT ASSIGNEE(S):

Byk Gulden Lomberg Chemische Fabrik Gmbh, Germany

PCT Int. Appl., 26 pp.

SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

1

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PA	TENT	KIND DATE			APPLICATION NO.						DATE						
Wo	2001	0727	55		A1 20011004			WO 2001-EP3510						- 2	 0010	 328	
	W:	ΑE,	AL,	ΑU,	BA,	BG,	BR,	CA,	CN,	CO	, CZ,	EE,	GE,	HR,	HU,	ID,	IL,
		IN,	JP,	KR,	LT,	LV,	MK,	MX,	NO,	ΝZ	, PL,	RO,	SG,	SI,	SK,	UA,	US.
		VN,	YU,	ZA,	ZW,	AM,	ΑZ,	BY,	KG,	ΚZ	, MD,	RU,	TJ,	TM	•		•
	RW:	AT,	ΒE,	CH,	CY,	DE,	DK,	ES,	FI,	FR	, GB,	GR,	IE,	IT,	LU,	MC,	NL,
		PT,	SE,	TR											-	•	
BR	2001	0095	39		Α		2003	0204		BR .	2001-	9589			2	0010	328
EP	1286				A1		2003	0305		EP :	2001-	9294	63		2	0010	328
	R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR	, IT,	LI,	LU,	NL,	SE,	MC,	PT.
		ΙE,	SI,	LT,	LV,	FI,	RO,	MK,	CY,	AL	, TR	•	•	•	•	•	,
	2003				Т2		2003			JP :	2001-	5706	64		2	0010	328
ИО	2002	0045	72		Α		2002	0924		NO :	2002-	4572				0020	924
US	2003	1005	78		A1		2003	0529		US :	2002-	1826	20		2	0021	001
PRIORITY	APP:	LN.]	INFO.	. :						EP :	2000-	1066	90	7	A 2	0000	329
											2001-					0010	
OTHER SO						PAT	135:	27296						·			

AB Compds. of formula I [R1 = Me, hydroxymethyl; R2-R5 = H, OH, OMe, OEt, OPr, OPr-i, OBu, methoxyethoxy, methoxypropoxy], are suitable for the prevention and treatment of gastrointestinal diseases. Thus, II is prepared and is shown to inhibit acid secretion 100% in rat stomach at 1 μ mol/kg.

IT 362605-90-1P 362605-91-2P 362605-92-3P 362605-93-4P 362605-94-5P 362605-96-7P 362605-97-8P 362605-98-9P 362605-99-0P 362606-00-6P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of pyrano[2,3-c]imidazo[-1,2-a]pyridine derivs. for treatment of gastrointestinal disorders)

RN 362605-90-1 CAPLUS

CN 7H-Imidazo[1,2-a]pyrano[2,3-c]pyridin-8-ol, 8,9-dihydro-7-(2-methoxyethoxy)-2,3-dimethyl-9-phenyl-, (7R,8S,9R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 362605-91-2 CAPLUS

CN 7H-Imidazo[1,2-a]pyrano[2,3-c]pyridin-8-ol, 8,9-dihydro-7-(2-methoxyethoxy)-2,3-dimethyl-9-phenyl-, (7S,8S,9R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

REFERENCE COUNT:

3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 6 OF 7 CAPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER:

2001:730747 CAPLUS

DOCUMENT NUMBER:

135:272962

TITLE: INVENTOR(S):

Preparation of alkylated imidazopyridine derivatives Postius, Stefan; Kromer, Wolfgang; Senn-Bilfinger,

Joerg; Buhr, Wilm

PATENT ASSIGNEE(S):

BYK Gulden Lomberg Chemische Fabrik GmbH, Germany;

Simon, Wolfgang-Alexander; Altana Pharma AG

SOURCE: PCT Int. Appl., 57 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND DATE	APPLICATION NO.	DATE			
WO 2001072754 WO 2001072754 WO 2001072754	C1 20030213	3	20010328			
W: AE, AL, AU, IN, JP, KR, VN, YU, ZA, RW: AT, BE, CH,	BA, BG, BR, CA, LT, LV, MK, MX, ZW, AM, AZ, BY,	CN, CO, CZ, EE, GE, HI NO, NZ, PL, RO, SG, SI KG, KZ, MD, RU, TJ, TN FI, FR, GB, GR, IE, II	I, SK, UA, US, M			
PT, SE, TR AU 2001044225 EP 1313739 R: AT, BE, CH, IE, SI, LT, BR 2001009542 JP 2003528876	A5 20011008 A1 20030528 DE, DK, ES, FR, LV, FI, RO, MK, A 20030610 T2 20030930 A 20030404	AU 2001-44225 EP 2001-917121 GB, GR, IT, LI, LU, NI CY, AL, TR BR 2001-9542 JP 2001-570663 ZA 2002-7636	20010328 20010328 L, SE, MC, PT, 20010328 20010328 20020923			
US 2003158193 PRIORITY APPLN. INFO.:		NO 2002-4597 US 2002-240039 EP 2000-106696 WO 2001-EP3507	20020925 20020927 A 20000329 W 20010328			

$$R^3$$
 R^4
 R^4 ?
 R^5 ?
 R^5
 R^5
 R^7
 $R^$

The title compds. I (R = H, alkyl, alkoxyalkyl, hydroxyalkyl; R2 = H, AΒ alkyl, hydroxyalkyl, halo, alkenyl, alkynyl; R3 = H, halo, F3C, alkyl, alkenyl, alkynyl, hydroxyalkyl, alkoxyalkyl carbamoyl; one of R4 and R4a is H, alkyl, alkenyl, Ph and the other is HO, alkoxy, alkoxyalkoxy, alkylcarbonyloxy, R4R4a = O, alkylidene; one of R5 and R5a is H, alkyl, alkenyl, Ph and the other is H, HO, alkoxy, alkoxyalkoxy, alkylcarbonyloxy, R5R5a = O, alkylidene; R6 = H, halo, alkyl, alkoxy, alkoxycarbonylamino, F3C; R7 = H, halo, alkyl, alkoxy; X = O, NH) were prepared for the prevention and treatment of gastrointestinal diseases. Thus, (8R9R)-2,3-dimethyl-8-hydroxy-9-phenyl-7,8,9,10tetrahydroimidazo[1,2-h][1,7]napnthyridin-7-one was methylated with MeI followed by reduction with NaBH4 to give (7R,8R,9R)-2,3,8-trimethyl-7,8dihydroxy-9-phenyl-7,8,9,10-tetrahydroimidazo[1,2-h][1,7]napnthyridine (II). At 1 μ mol/kg (i.v.) II inhibited acid secretion of the perfused rat stomach stimulated pentagastrin by 100%.

IT 364041-33-8P 364041-34-9P 364041-35-0P 364041-36-1P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of alkylated imidazopyridine derivs.)

RN 364041-33-8 CAPLUS

CN 7H-Imidazo[1,2-a]pyrano[2,3-c]pyridine-7,8-diol, 8,9-dihydro-2,3,7-trimethyl-9-phenyl-, (7S,8R,9R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 364041-34-9 CAPLUS

CN 7H-Imidazo[1,2-a]pyrano[2,3-c]pyridine-7,8-diol, 8,9-dihydro-2,3,7-trimethyl-9-phenyl-, (7R,8R,9R)- (9CI) (CA INDEX NAME)

REFERENCE COUNT:

THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 7 OF 7 CAPLUS COPYRIGHT 2004 ACS on STN L4

3

ACCESSION NUMBER:

1998:795020 CAPLUS

DOCUMENT NUMBER:

130:25073

TITLE:

Preparation of fused dihydropyrans for use in the prevention and treatment of gastrointestinal diseases Grundler, Gerhard; Simon, Wolfgang-alexander; Postius,

INVENTOR(S):

Stefan; Riedel, Richard; Thibaut, Ulrich;

Senn-bilfinger, Jorg

PATENT ASSIGNEE(S):

Byk Gulden Lomberg Chemische Fabrik Gmbh, Germany

SOURCE:

PCT Int. Appl., 33 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PA	TENT	NO.			KIND DATE			APPLICATION NO.							DATE			
WO		AL, LV, AM,	AU, MK, AZ, BE,	BA, MX, BY,	BG, NO, KG,	BR, NZ, KZ,	1998 CA, PL, MD, DK,	CN, RO, RU,	CZ, SG, TJ,	EE SI TM	i, (GE, SK,	HU, TR,	ID, UA,	IL, US,	JP, VN,	KR, YU,	LT, ZW,
AU	9879	154			A1		1998	1230		AU	199	98-7	9154	1		1	9980	523
AU	7367	67			B2		2001	0802						-				223
EP	9849	59			A1		2000	0315		ΕP	199	98-9	2937	70		1	9980	523
	R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR	, 1	ľΤ,	LI,	LU.	NL.	SE.	MC.	PT.
		ΙE,	SI,	LT,	LV,	FI,	RO				·	•	•			,	,	,
	98091	185			A		2000	0801		BR	199	98-9	185			1	9980	523
JP	20015	52670)3		T 2		2001	1218									9805	
ZA	98044	163			A		1998	1130									9805	
US	61601	L19					20003		Ţ								9911	
	99109				A		20000	0430									9991	125
PRIORITY	APPI (LN. 1	NFO.	:										4			9705	
														7		_	9805	
OTHER SC	URCE	(S):			MARP	ΆΤ	130:2	25073							•			

$$R^3$$
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AB Fused dihydropyrans I [R1 = alkyl; R2 = alkyl, hydroxyalkyl; R3 = H, halogen; R4 = R5 = H, OH, alkoxy, alkylcarbonyloxy, oxo; R4R5 = fused heterocycle, such as OCH2O or O(CH2)2O; R6 = H, CF3, halogen, alkoxy, alkoxycarbonylamino; R7 = H, halogen, alkyl, alkoxy] were prepared for the prevention and treatment of gastrointestinal diseases. Thus, cis-I [R1 = R2 = Me, R3 = R4 = R6 = R7 = H, R5 = OH] was prepared by reaction of II with Bu3SnH/AIBN in benzene followed by treatment with saturated KOH solution The prepared compds were tested for inhibition of acid secretion on perfused rat stomach.

IT 216159-49-8P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of fused dihydropyrans for use in the prevention and treatment of gastrointestinal diseases)

II

RN 216159-49-8 CAPLUS

CN 7H-Imidazo[1,2-a]pyrano[2,3-c]pyridine-7,8-diol, 8,9-dihydro-2,3-dimethyl-9-phenyl-(9CI) (CA INDEX NAME)

REFERENCE COUNT:

THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

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(FILE 'HOME' ENTERED AT 10:17:19 ON 04 SEP 2004)

FILE 'REGISTRY' ENTERED AT 10:17:36 ON 04 SEP 2004

L1 STRUCTURE UPLOADED

L2 2 S L1

L3 33 S L1 FULL

FILE 'CAPLUS' ENTERED AT 10:18:06 ON 04 SEP 2004

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L4

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L1 HAS NO ANSWERS

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Structure attributes must be viewed using STN Express query preparation.

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PALM INTRANET

Day: Saturday Date: 9/4/2004 Time: 09:40:25

Inventor Name Search Result

Your Search was:

Last Name = SENN-BILFINGER

First Name = J

Application#	# Patent#	Status	Date Filed	Title	Inventor
10851092	Not Issued	020	05/24/2004	POLYSUBSTITUTED IMIDAZOPYRIDINES	SENN-BI JORG
10826337	Not Issued	020	04/19/2004	PRODRUGS OF IMIDAZOPYRIDINE DERIVATIVES	SENN-BI JORG
10811496	Not Issued	030	04/01/2004	PYRANO[2,3-C]IMIDAZO[-1,2-A]PYRIDINE DERIVATIVES FOR THE TREATMENT OF GASTROINTESTINAL DISORDERS	SENN-BI JORG
10783512	Not Issued	030	02/23/2004	TETRAHYDROPYRIDOETHERS	SENN-BI JORG
10667524	Not Issued	092	09/23/2003	PROCESS AND INTERMEDIATES FOR THE PREPARATION OF IMIDAZOPYRIDINES	SENN-BI JORG
10485514	Not Issued	020	02/02/2004	ALKYL-SUBSTITUTED IMIDAZOPYRIDINES FOR THE TREATMENT OF GASTROINTESTINAL DISORDERS	SENN-BI JORG
10485512	Not Issued	020		AMINO-SUBSTITUTED IMIDAZOPYRIDINES FOR THE TREATMENT OF GASTROINTESTIAL DISEASES	SENN-BI JORG
<u>10485418</u>	Not Issued	030	01/30/2004	TRICYCLIC EPOXIDES	SENN-BI JORG
10482483	Not Issued	020		PROCESS FOR THE PRODUCTION OF 3-PHENYLISOSERINE	SENN-BI JORG
<u>10380624</u>	Not Issued	094		·	SENN-BI JORG
10240039	Not Issued	061	6.8	TATED IX I A PRIZE AND G	SENN-BI JVRG
10182654	<u>6696461</u>	150	10/04/2002	:	SENN-BI JVRG
10182652	<u>6653477</u>	150	09/19/2002		SENN-BI JORG

10182619	Not Issued	161	10/01/2002	PRODRUGS OF IMIDAZOPYRIDINE DERIVATIVES	SENN-BI JVRG
10149290	6716990	150	06/11/2002	PROCESS AND INTERMEDIATES FOR THE PREPARATION OF IMIDAZOPYRIDINES	SENN-BI JORG
10103733	6696460		03/25/2002	TETRAHYDROPYRIDOETHERS	SENN-BI JORG
09926267	6503923	150	10/03/2001	HALOALKOXY IMIDAZONAPHTHYRIDINES	SENN-BI JORG
09807970	6384048	150	04/27/2001	IMIDAZONAPHTHYRIDINES	SENN-BI JORG
09582212	6436953	150	07/19/2000	TETRAHYDROPYRIDOETHERS	SENN-BI JORG
09423626	6160119	150	11/16/1999	FUSED DIHYDROPYRANS	SENN-BI JORG
<u>09381617</u>	<u>6197783</u>	150	09/24/1999	TETRAHYDROPYRIDO COMPOUNDS	SENN-BI , JORG
09117139	6096758	150	07/24/1998	3-METHYLIMIDAZOPYRIDINES	SENN-BI , JORG
<u>08776391</u>	Not Issued	161	04/17/1997	ACYLIMIDAZOPYRIDINES	SENN-BI , JORG
<u>08776390</u>	6124313	150	05/16/1997	IMIDAZOPYRIDINE AZOLIDINONES	SENN-BI , JORG
08776349	Not Issued	161	05/05/1997	BENZYLIMIDAZOPYRIDINES	SENN-BI , JORG
08776348	Not Issued	161	01/28/1997	HALOIMIDAZOPYRIDINES	SENN-BI , JORG
<u>08776047</u>	6162809	150		THIOPYRIDYL COMPOUNDS FOR CONTROLLING HELICOBACTER BACTERIA	SENN-BI , JORG
08765980	5922720	150		PIPERAZINE THIOPYRIDINES FOR THE CONTROL OF HELICOBACTER BACTERIA	SENN-BI , JORG
08750792	6107312	150		THIOPYRIDINES FOR USE IN THE CONTROL OF HELICOBACTER BACTERIA	SENN-BI , JORG
08750785	5859030	150]	SUBSTITUTED ARYLALKYLTHIOALKYLTHIOPYRIDINES FOR USE IN THE CONTROL OF HELICOBACTER BACTERIA	SENN-BI , JORG
08652505	5668131	150	1 51	SUBSTITUTED AMINOALKYLAMINOPYRIDINES	SENN-BI , JORG
08537772	<u>5824687</u>	150	10/20/1995	PYRIDINIUM SALTS AND THEIR USE FOR	¥

				BACTERIA	
08505271	<u>5665730</u>	150	08/15/1995	PHARMACEUTICALLY USEFUL IMIDAZOPYRIDINES	SENN-BI , JORG
08505270	5587389	150	08/15/1995	SUBSTITUTED HETEROARYLAKLYLTHIOPYRIDINES FOR CONTROLLING HELICOBACTER BACTERIA	SENN-BI , JORG
08295681	Not Issued	161	08/25/1994	NOVEL AND KNOWN DISULFIDES AND THEIR USE IN THE CONTROL OF HELICOBACTER BACTERIA	SENN-BI , JORG
07445611	5112834	150	01/16/1990	IMIDAZOLE PROTECTORANT FOR THE STOMACH AND INTESTINE	SENN-BI , JORG
06437883	4472409	250	10/29/1982	2-PYRIDYLMETHYL THIO(SULFINYL)BENZIMIDAZOLES WITH GASTRIC ACID SECRETION INHIBITING EFFECTS	SENN-BI , JORG
<u>06344172</u>	<u>4363816</u>	250		TRICYCLIC PYRROLES, THEIR COMPOSITIONS AND THEIR USE	SENN-BI , JORG

Inventor Search Completed: No Records to Display.

	Last Name	First Name	
Search Another:	Senn-Bilfinger	J	
Inventor		Search	······

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